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RESEARCH ARTICLE

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# Intragenic homogenization and multiple copies of prey-wrapping silk genes in *Argiope* garden spiders

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## Abstract

**Background:** Spider silks are spectacular examples of phenotypic diversity arising from adaptive molecular evolution. An individual spider can produce an array of specialized silks, with the majority of constituent silk proteins encoded by members of the spidroin gene family. Spidroins are dominated by tandem repeats flanked by short, non-repetitive N- and C-terminal coding regions. The remarkable mechanical properties of spider silks have been largely attributed to the repeat sequences. However, the molecular evolutionary processes acting on spidroin terminal and repetitive regions remain unclear due to a paucity of complete gene sequences and sampling of genetic variation among individuals. To better understand spider silk evolution, we characterize a complete aciniform spidroin gene from an *Argiope* orb-weaving spider and survey aciniform gene fragments from congeneric individuals.

**Results:** We present the complete aciniform spidroin (*AcSp1*) gene from the silver garden spider *Argiope argentata* (*Aar\_AcSp1*), and document multiple *AcSp1* loci in individual genomes of *A. argentata* and the congeneric *A. trifasciata* and *A. aurantia*. We find that *Aar\_AcSp1* repeats have >98% pairwise nucleotide identity. By comparing *AcSp1* repeat amino acid sequences between *Argiope* species and with other genera, we identify regions of conservation over vast amounts of evolutionary time. Through a PCR survey of individual *A. argentata*, *A. trifasciata*, and *A. aurantia* genomes, we ascertain that *AcSp1* repeats show limited variation between species whereas terminal regions are more divergent. We also find that average dN/dS across codons in the N-terminal, repetitive, and C-terminal encoding regions indicate purifying selection that is strongest in the N-terminal region.

**Conclusions:** Using the complete *A. argentata AcSp1* gene and spidroin genetic variation between individuals, this study clarifies some of the molecular evolutionary processes underlying the spectacular mechanical attributes of aciniform silk. It is likely that intragenic concerted evolution and functional constraints on *A. argentata AcSp1* repeats result in extreme repeat homogeneity. The maintenance of multiple *AcSp1* encoding loci in *Argiope* genomes supports the hypothesis that *Argiope* spiders require rapid and efficient protein production to support their prolific use of aciniform silk for prey-wrapping and web-decorating. In addition, multiple gene copies may represent the early stages of spidroin diversification.

**Keywords:** *AcSp1*, Concerted evolution, Intragenic homogenization, Multiple loci, Silk, Spidroin, Spider

## Background

Spider silks are ideal for studying the molecular evolutionary processes that create and maintain adaptive characteristics. An individual spider can produce and use different silk types singly or in combination for specific tasks, with each silk type having mechanical properties well-suited to its function. For example, aciniform silk is used in prey immobilization and egg sac construction [1,2]. The

mechanical properties of aciniform silk include impressive extensibility and toughness [1], making it excellent for swathing struggling prey because it is easy to stretch but difficult to break. Orb-weaving garden spiders from the genus *Argiope* are renowned for their use of aciniform silk. *Argiope* employ many layers of aciniform silk to completely immobilize and envelop their prey (e.g. [3,4]), and *Argiope* are also a model system for studying the purpose of aciniform-silk web decorations, known as stabilimenta, that have been implicated in predator avoidance, prey attraction, and web stability (for review see [5,6]).

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Of the five fibrous silks spun by the silver garden orb-weaver *Argiope argentata*, aciniform silk is the toughest and one of the most extensible [7]. However, little is known about the evolution of aciniform silk's physical attributes. Spider silk mechanical properties are related to the suite of proteins that compose each silk type. The majority of spider silk proteins, or spidroins (a contraction of "spider-fibroins" [8]), are encoded by members of a single gene family. Spidroins are typically very large (>200 kDa), and are dominated by a series of iterated repeats flanked by short amino (N)- and carboxy (C)-terminal regions [9,10]. The length, number, and amino acid (aa) composition of the iterated repeats are silk-type specific, whereas phylogenetic analyses have shown that aa residues in the N- and C-terminal regions are more conserved across spidroins [11,12]. Repeat aa sequence corresponds to secondary structures that are partly responsible for silk mechanical properties (e.g. [13-15]), and conservation of the N- and C-terminal regions [12] and their presence in spun silk fibers suggests an important role in spider silk biology [16-18].

The evolutionary maintenance of spidroin repeat sequences within a silk type and the divergence of those repeat sequences between silk types is central to spider silk function and diversity. Within a particular spidroin, repeat units tend to be highly similar, or homogeneous, in amino acid and nucleotide sequence. The gene encoding aciniform spidroin (*AcSp1*) has repeats that are relatively complex among spidroin family members, however, despite this complexity, *AcSp1* repeats are also spectacularly homogenized [1,19]. A recent analysis of a complete *AcSp1* from the western black widow *Latrodectus hesperus* showed that its repetitive region, like those of other spidroins, is dominated by the amino acids glycine (G), alanine (A), and serine (S) [19]. However, *L. hesperus* *AcSp1* repeats have few or none of the short glycine and alanine-rich subunits, such as GGX, poly-GA, and poly-A, that can be the bulk of other spidroin repeats [9]. Nevertheless, *L. hesperus* *AcSp1* repeats are remarkably homogenized (>99% identity at the nucleotide level [19]). This is consistent with results from a partial length *AcSp1* cDNA from the banded garden orb-weaver *Argiope trifasciata*, which has 14 repeats that are each 600 bp and share 99.9% identity at the nucleotide level [1].

The high level of *AcSp1* repeat homogeneity is frequently attributed to gene conversion or unequal crossing over resulting in intragenic concerted evolution (e.g. [1,19,20]). Concerted evolution usually refers to homogenization among gene family members, such as rDNA gene copies [21], but it can also occur within a gene [22,23]. Stabilizing selection alone would maintain protein sequence, resulting in a high level of repeat identity at the aa level. However, the extreme level of homogenization reported for *AcSp1* repeats provides evidence for concerted

evolution because it exists at both the protein and nucleotide levels [1,19].

In addition to concerted evolution, repeat homogeneity in *AcSp1* may be maintained by functional constraints. Recent nuclear magnetic resonance (NMR) studies of *AcSp1* repeats from both *Nephila antipodiana* and *A. trifasciata* delineate different domains in each repeat unit, one domain that is rich in alpha helices and one that is not [24,25]. Xu et al. [25] used NMR and dihedral angles from global likelihood estimate (DANGLE) analyses to predict the chemical shift indices of a 199 aa recombinant *A. trifasciata* *AcSp1* repeat. The consensus secondary structure assignments specified that the last quarter of the protein was unstructured, but that the first three-quarters of the repeat contained six major helical regions. Protein structures such as these six alpha helices are considered the foundation for silk mechanical properties (e.g. [25,26]).

Assessing the extent to which a spidroin is homogenized within a single gene or among individuals is difficult because the repetitive region makes it exceptionally challenging to sequence complete spidroin genes. Indeed, partial length sequences that are biased toward the C-terminus greatly dominate the number of published spidroins [12]. Additionally, the evolutionary processes leading to spidroin divergence between species and silk types are often unclear due to a lack of knowledge about spidroin genetic variation among individuals.

Here, we address these issues by presenting a complete spidroin gene from an *Argiope* spider, the *AcSp1* sequence of *A. argentata* (*Aar\_AcSp1*), and by screening for *AcSp1* variation among individual *A. argentata*, *A. trifasciata*, and *A. aurantia* spider genomes. Sequencing the full array of *Aar\_AcSp1* repeats enabled us to test hypotheses of concerted evolution and functional constraints. Based on previous spidroin research, *Aar\_AcSp1* repeats should be extremely homogenous at the nucleotide and amino acid levels. In addition, amino acid sequences that are predicted to correspond to the structural motifs that contribute to the toughness and extensibility of aciniform silk should be more conserved between *Argiope* species relative to surrounding regions. Among the surveyed *A. argentata* individuals, we expected *Aar\_AcSp1* to be a single-copy gene, similar to *L. hesperus* *AcSp1* [19]. Between species, previous research suggests that the spidroin repeats within each silk type are highly conserved, but that the terminal regions show more variation [12], and we hypothesized that *Aar\_AcSp1* would also follow this pattern.

## Results and discussion

### *Argiope argentata* *AcSp1* complete sequence and phylogenetic placement

Despite obtaining 59 *AcSp1* cDNA clones, including one that was >8 kb [1], a complete *Argiope* *AcSp1* remained

elusive until the present study. By screening a large-insert genomic DNA library, we sequenced and assembled 18,080 bp of *A. argentata* DNA including a complete open reading frame that is 13,440 bp long and predicted to encode a 4,479 aa *A. argentata* AcSp1 (*Aar\_AcSp1*; Figure 1; GenBank KJ206620). No introns were detected. The putative protein has a predicted size of ~430 kDa, and the most abundant amino acids are serine (22.6%), alanine (14.4%) and glycine (13.3%). *Aar\_AcSp1* has three regions, a central repetitive region that is flanked by conserved N- and C-terminal regions. The repetitive region dominates ~90% of the protein and is composed of 20 iterated repeats (Figure 1). The first 19 repeats are each 204 aa, and the last repeat is 186 aa due to truncation at the end (Figure 1). The length, amino acid composition, and organization of AcSp1 are all consistent with other spidroin family members [9].

Phylogenetic analyses of *Aar\_AcSp1* N- and C-terminal coding regions with those from other spidroins grouped *Aar\_AcSp1* with the *Latrodectus* (widow spider) AcSp1 sequences in a well-supported clade (bootstrap value = 98%; Figure 2). *Latrodectus* and *Argiope* are estimated to have diverged from each other ~175 million years ago (MYA) [27]. Despite this lengthy time period, the recovery of an AcSp1 clade was consistent with prior studies in which spidroin sequences nearly always grouped based on silk type (e.g. [9,10]). The sister group to the AcSp1 clade was TuSp1, tubuliform (egg-case) spidroin, suggesting that these paralogs have a relatively recent common ancestor [19]. Further potential evidence of their shared ancestry is that both of these silk types are used in egg-case construction and both have repeats that are relatively complex compared to other spidroins [28]. In our phylogenetic analysis, a large, weakly supported assemblage of spidroins is sister to the combined AcSp1 and TuSp1 clade. Given the low support, it is unclear which spidroins are most closely related to AcSp1 and TuSp1.

#### ***Argiope argentata* AcSp1 repeat homogeneity**

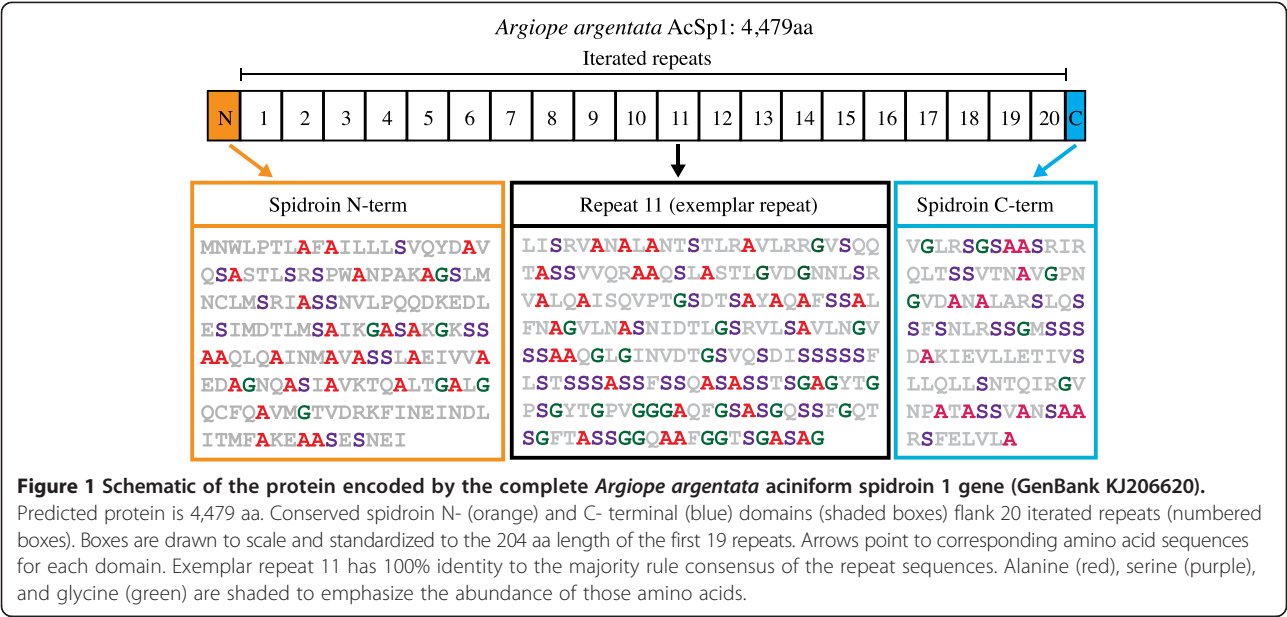
As expected, *Aar\_AcSp1* repeats are complex and spectacularly homogenized. Although glycine, alanine, and serine account for ~50% of its repetitive region composition, *Aar\_AcSp1* has few of the glycine/alanine-rich motifs such as GGX, GPG, poly-GA, and poly-A that are dominant in the dragline major ampullate spidroins (MaSp1, MaSp2) from *Argiope* and other taxa [9]. At the nucleotide level, the average pairwise percent identity between *Aar\_AcSp1* repeats is an astonishing 98.7%. Complexity and extreme homogenization are also features of previously described AcSp1 sequences [1,19].

The extreme nucleotide identity of *Aar\_AcSp1* is consistent with concerted evolution, and cannot be easily explained by codon usage bias. For example, *Aar\_AcSp1* codon use is strongly influenced by amino acid position

within a repeat. In our repeat alignment (Additional file 1: Figure S1), the neighboring alanine codons at nucleotide positions 103–105 and 106–108 are GCC and GCT, respectively. GCCGCT is present in the same relative location in all twenty repeats. Similarly, the glycine codons that appear at nucleotide positions 64–66 and 130–132 also consistently use different codons (GGT and GGA, respectively). The same alternative codons are used at the same exact positions throughout most, if not all, the repeats. Despite a slight skew toward alanine codons that end in adenine (A) or thymine (T) (55.0% GCW, W being the IUPAC ambiguity code for A or T; Additional file 2: Table S5), it is difficult to postulate that selective forces acting at the level of codon usage are responsible for the extensive homogeneity of codon positions found throughout the 612 bp *Aar\_AcSp1* repeat. Concerted evolution that fixes particular codons at particular locations across repeats provides a clearer explanation.

Analyses of the full array of *Aar\_AcSp1* iterated repeats were also consistent with two concerted evolution predictions. First, ML analysis grouped araneid AcSp1 repeats into well-supported, species-specific clades rather than grouping the repeats across species (Figure 3A). Furthermore, nucleotide pairwise identity within each species averaged 98%, but pairwise identity between *Aar\_AcSp1* repeats and repeats from other species averaged only 78.5% (73.6% vs. *Araneus ventricosus*, 79.1% vs. *A. trifasciata*, and 82.8% vs. *A. amoena*). That repeats are more similar within species than between species regardless of intragenic repeat position can be explained by rapid intra-specific spread of genetic variation via unequal crossing over during recombination [22,23].

Second, the average nucleotide pairwise identity of the first and last *Aar\_AcSp1* repeats to the rest of the array is slightly lower at 96% and 93%, respectively. Less similar first and last repeats are consistent with some models of concerted evolution [29]. However, araneid AcSp1 first and last repeats still grouped within species-specific clades (Figure 3A), suggesting that these repeat sequences are more homogeneous within a gene than those of previously analyzed spidroins. For example, in an analysis of repetitive units from the flagelliform spidroin (*Flag*) of the golden orb-weaver *Nephila clavipes* and the congeneric *Nephila inaurata madagascariensis*, the first repeats grouped together across species, and the last repeats also formed their own clade. By contrast, the central (not first or last) repeats formed species-specific clades because each repetitive unit was nearly identical within each species yet divergent across species [30]. Longer estimated divergence times between the species in our present study may explain the more thorough homogenization of araneid AcSp1 sequences compared to that of the previously studied *Nephila Flag* sequences. The estimated divergence time between the *Nephila* species is ~7.4 MYA [31],



whereas *Araneus* and *Argiope* are thought to have diverged ~30 MYA and within *Argiope*, ~23 MYA between *A. argentata* and *A. trifasciata* [32].

**Functional constraints on AcSp1 repeats**

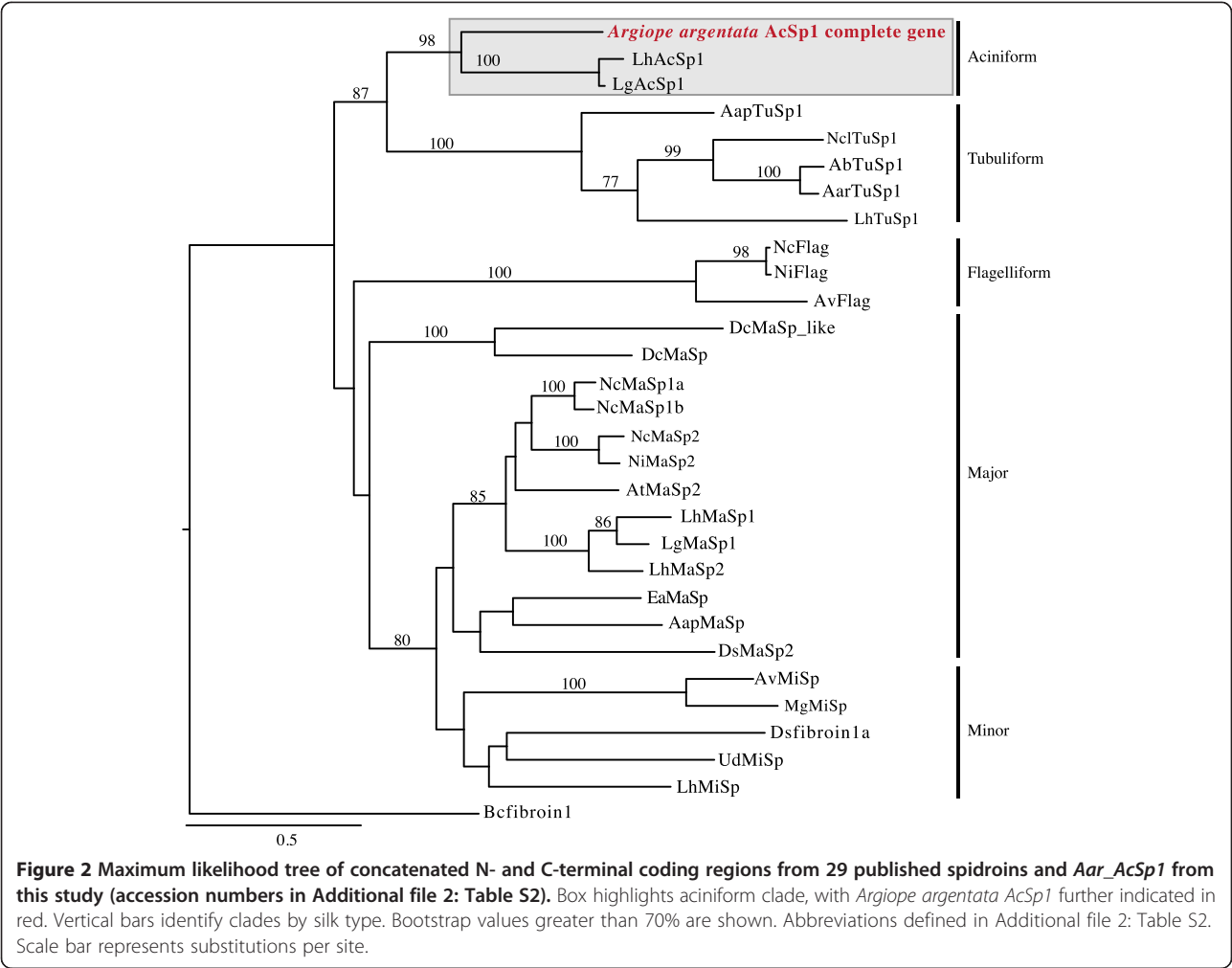
We predicted that functional constraints would result in greater aa sequence conservation in the portion of each repeat proposed by Xu et al. to contain six alpha helices [25]. To test this, we first compared known Araneidae repetitive regions. We aligned consensus AcSp1 repeat sequences from three *Argiope* species (*A. argentata*, *A. trifasciata*, *A. amoena*) and *Araneus ventricosus*. We then graphed pairwise identities for each aligned position between the *A. trifasciata* repeat sequence and the other species, and plotted it against the predicted *A. trifasciata* domains from Xu et al. [25] (Figure 3B). Using amino acid positions from Xu et al. [25], the average percent pairwise identity over the 150 aa helix-rich domain was 84.0%, but only 54.1% over the remaining 49 aa. Xu et al. [25] also noted a major alpha-helical domain from 102–151 aa, encompassing the region denoted as helix 5 and 6 in Figure 3B. Consistent with being structurally important, the average percent identity in this domain was 90.7%. Moreover, our alignment was slightly longer (216 aa) than the *A. trifasciata* recombinant repeat length (199 aa) due to indels that only appeared in the unstructured region. Notably, in the region from 200–209 aa (our alignment), the *A. trifasciata* repeat has a deletion (Figure 3B). These indels further indicate that the final quarter of AcSp1 repeats is less conserved than the first three-quarters.

To investigate amino acid conservation in the predicted AcSp1 helical regions across greater evolutionary time, we also aligned consensus amino acid AcSp1 repeat sequences from *L. hesperus* and *Uloborus diversus* to the *A. trifasciata* repeat from Xu et al. [19,25]. Araneidae, represented here by *Argiope* and *Araneus*, and Theridiidae, represented by *L. hesperus*, are members of the superfamily Araneoidea, with araneids and theridiids estimated to have last shared a common ancestor ~175 MYA [27]. *U. diversus* is within the Deinopoidea, the sister-group to the Araneoidea. Araneoids and deinopoids diverged from each other ~210 MYA [27]. Together, Araneoidea and Deinopoidea compose the Orbiculariae (orb-web weaving spiders and their relatives).

The AcSp1 repeat units from *L. hesperus* and *U. diversus* are almost twice as long as the araneid AcSp1 repeat units. The *L. hesperus* and *U. diversus* repeat units can be further subdivided into two parts that align with each other [19]. We aligned each part from each species (two parts per species) to the *A. trifasciata* repeat separately. We then calculated the average pairwise percent identity for each comparison and for each of the six putative alpha-helical regions predicted by Xu et al. ([25]; Additional file 2: Table S6). The overall pairwise identities between *L. hesperus* repeat part 1 and *U. diversus* repeat part 1 with the *A. trifasciata* repeat was 30% and 29%, respectively. Of note, the percent pairwise identity between *L. hesperus* repeat part 1 and the *A. trifasciata* repeat was 47% in the *A. trifasciata* region associated with helix 4, and it was 41% against *U. diversus* repeat part 1 in the region associated with helix 6. 47% and 41% were the highest pairwise identity percentages.

Our results strongly support the hypothesis that functional constraints are acting to conserve protein sequence in the repetitive region of AcSp1. Our comparison of *A. trifasciata* AcSp1 repeat sequence with that of other





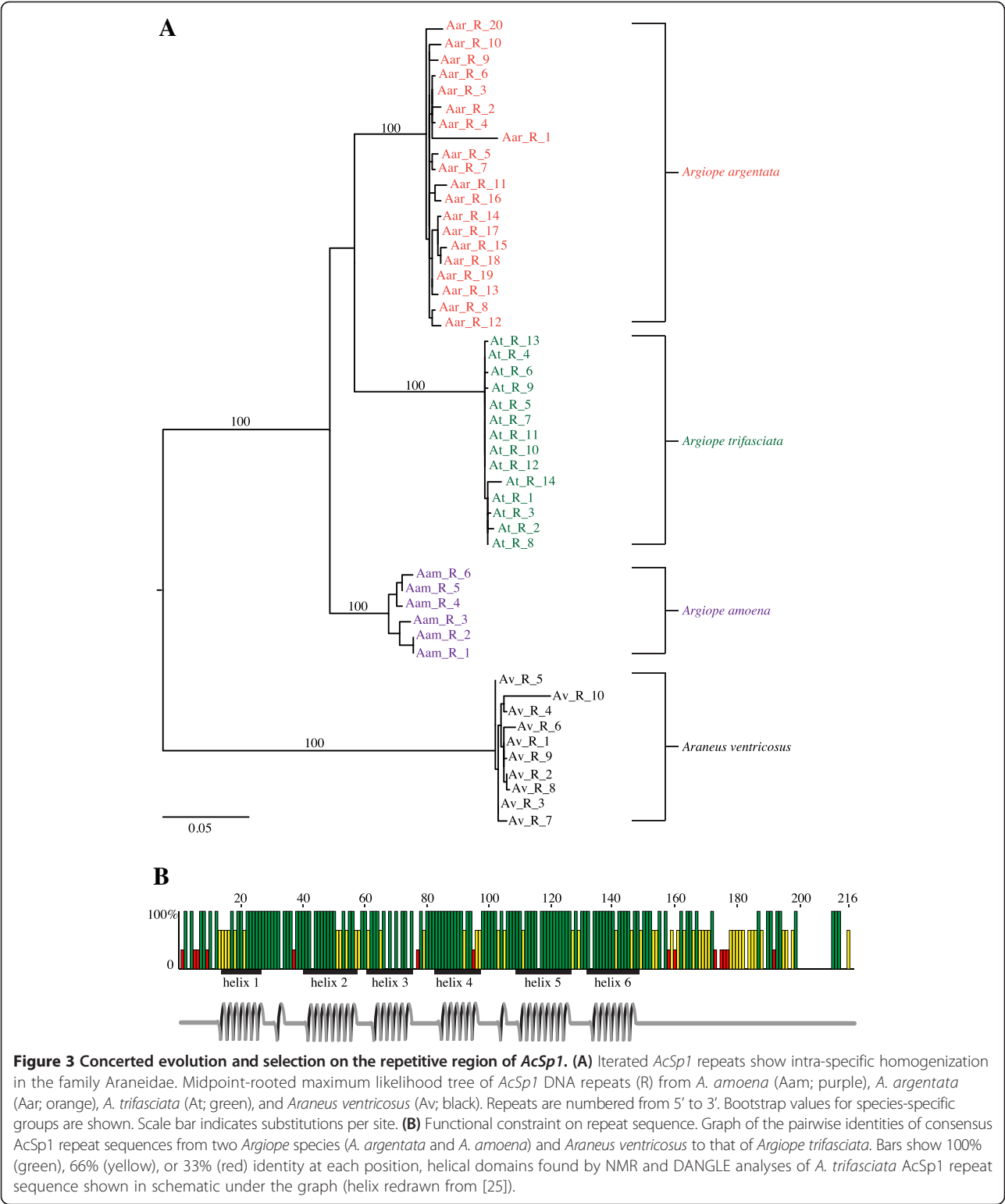
araneid species indicates that a specific amino acid sequence is maintained in the predicted helix-rich domain of *AcSp1* repeats across Araneidae. In contrast, comparison between the *A. trifasciata* repeat with part 1 of repeats from *L. hesperus* and *U. diversus* indicates that amino acid sequence in the regions associated with alpha-helices 4 and 6 are the most highly conserved across Orbiculariae. The higher level of conservation in the amino acid sequences corresponding to helices 4 and 6 may indicate that these regions impart the same general function across Orbiculariae whereas the other predicted helical regions of *A. trifasciata* impart functions unique to Araneidae. Sequencing *AcSp1* from other genera of Araneidae and other families of Orbiculariae will enable further elucidation of these hypotheses.

To our knowledge, there are no current predictions about the secondary structures of *L. hesperus* or *U. diversus* *AcSp1* repeats. It is feasible that, like the *AcSp1* domains of *N. antipodiana* and *A. trifasciata*, *L. hesperus* and *U. diversus* *AcSp1* repeats also feature distinct structural regions. Finally, our analysis may be an underestimation

of sequence conservation because it does not include amino acid replacements that are functionally equivalent. However, predicting functional protein similarity is difficult given the extensive physicochemical changes that spider silk undergoes as it is processed from a liquid into dry silk (e.g. [33,34]).

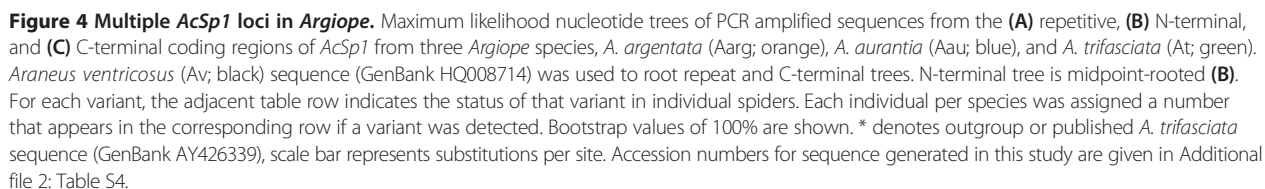
**Delineation of *AcSp1* variants in individual *Argiope* spiders**

Spidroin sequence variation between individual spiders is an important source of genetic variation for the evolution of different silk types within and between species. To investigate genetic variation in *AcSp1* between individuals of *A. argentata* and the congeneric *A. trifasciata* and *A. aurantia*, we first designed PCR primers targeting the repetitive region of *Aar\_AcSp1*. Amplification of genomic DNA across species and individuals resulted in *AcSp1* repeat sequences that did not show intraspecific variation but had significant inter-specific variation (Figure 4A). Intraspecific homogenization of the repeats could be explained by biased PCR amplification of a single repeat type in the repetitive region, however, our results are consistent



with the high degree of repeat sequence conservation in *AcSp1* sequences from araneids (Figure 3A) and *L. hesperus* [19].

Next, we designed PCR primers targeting N- and C-terminal coding regions of *Aar\_AcSp1*. We then amplified the same individual genomic DNAs that were surveyed





for the repetitive region (Figure 4). Unlike the repetitive region PCR, direct sequencing of terminal region PCR products resulted in extensive numbers of multiple peaks and in some cases, poor sequencing reads due to length differences. Thus, all terminal region PCR products were cloned, a total of 385 amplicons were sequenced, and *AcSp1* variants were diagnosed. Each variant was supported by at least two amplicons with sequences that had greater than 95% identical bases (Figure 4; see Methods). Unlike the repetitive region sequences, which showed no intra-specific or allelic variation, all terminal region amplifications were heterogeneous.

The number of variants characterized was surprising because all of the individual spiders surveyed were found to have more than two terminal region variants, indicating that these *Argiope* species must have multiple *AcSp1* encoding loci. *Argiope* spiders are not known to be polyploid, thus multiple gene copies per genome is the only explanation for more than two N- or C- terminal region variants in a single individual. For example, we found seven C-terminal variants in one *A. argentata*, suggesting at least four *AcSp1* gene copies (Figure 4C). Each *A. trifasciata* individual possessed a minimum of seven N- or C-terminal variants, again indicating at least four gene copies (Figure 4B, C). Likewise, an *A. aurantia* individual possessed six N-terminal variants but only two C-terminal variants (Figure 4B, C). The smaller number of C-terminal variants could be explained by lack of variation in the C-terminal region or by incomplete sampling of variants by PCR survey.

ML analysis of sequences from the PCR survey shows that the branch lengths in the repetitive region (Figure 4A) are shorter than the branch lengths of the terminal region trees (Figure 4B, C). The majority of N- and C-terminal variants cluster into well-supported, species-specific groups, and intra-specific branch lengths are very short compared to inter-specific branch lengths (Figure 4B, C). One exception is *A. argentata* C-terminal coding region variant V1, which forms a weakly supported group with *A. trifasciata* and *A. aurantia* C-terminal coding region variants (Figure 4C). Given the weak clade support, this variant is probably an outlier that is not as homogenized as the other *A. argentata* variants.

The shorter branch lengths of the repetitive region variants tree compared to those of the N- and C-terminal region trees suggest that the repetitive region is the most conserved araneid *AcSp1* region (Figure 4). Yet, comparison of the average ratio of non-synonymous to synonymous substitution rates (dN/dS) across codons implies that the N-terminal region has been subject to slightly stronger purifying selection than the repetitive and C-terminal regions (0.20 vs. 0.30 and 0.42 dN/dS, respectively). dN/dS estimations, however, assume independence of sites and thus are confounded by factors

such as concerted evolution and recombination. The full-length *Aar\_AcSp1* and other *AcSp1* provide extensive evidence that the repetitive region units are most likely not evolving independently from each other (Figure 3A; [1,19]). Thus concerted evolution and purifying selection both must play a role in the near-perfect homogeneity of *Argiope AcSp1* iterated repeats. Recombination can also affect tests of selection [35]. Because we could not conclusively determine the exact number of loci within an individual or assign alleles to specific loci, we were unable to ascertain recombination between loci. Subsequent analyses with additional data could address the impact of recombination on dN/dS estimates.

Previous work with *AcSp1* sequences did not find evidence for multiple loci [1,19]. The lack of variation among *A. trifasciata AcSp1* cDNA clones [1] may be due to over-expression or preferential cloning of one variant and thus its preponderance in the characterized cDNAs. Alternatively, consistent depletion of aciniform silk may be required to stimulate transcription of multiple *AcSp1* loci. This hypothesis is supported by a significant increase in aciniform-silk dependent web-decorating behavior in three species of *Argiope* in response to a two-week period of aciniform silk depletion [4]. Future work could focus on comparing the number of *AcSp1* variants expressed by spiders consistently depleted of aciniform silk versus that from spiders that are not depleted.

Survey of individual *L. hesperus* genomes also did not find *AcSp1* variants [19]. However, the detection of multiple *AcSp1* loci in *Argiope* but not *Latrodectus* is consistent with the hypothesis that *Argiope* spiders maintain multiple gene copies as a strategy for efficiently producing large amounts of protein. In contrast with *Argiope* spiders, *Latrodectus* spiders use markedly fewer strands of aciniform silk during prey capture [3] and do not make stabilimenta. Increased *AcSp1* copy number in *Argiope* spiders may therefore be a strategy for increasing protein production [36]. Because spidroins are costly, highly expressed proteins [37,38], resource abundance in the form of prey availability may also stimulate aciniform spidroin production in *Argiope* to prepare for resource scarcity [39,40]. Precedent for this strategy exists. In the bacteria *Escherichia coli*, multiple copies of rRNA operons provide a competitive advantage by enabling increased growth rates and decreased cell division lag time in environments where resources fluctuate rapidly [41,42].

Previous research has found variants for other spidroins [43-46], and that the dragline spidroin *MaSp1* is encoded by multiple loci in several species [47,48]. Unlike *A. argentata AcSp1* variants, the C-terminal coding region of *L. hesperus MaSp1* is nearly identical across loci [47]. This difference could indicate functional constraints on the C-terminal coding region of *MaSp1* that either differ from or are not acting on *AcSp1*. A comparison of

structural component predictions from the amino acid sequences of terminal regions across different spidroins would greatly inform our understanding of the contribution of the terminal regions to the evolution of different spider silk types.

## Conclusions

The highly similar iterated repeat array of our complete *Argiope argentata* *AcSp1* gene combined with sequence conservation of functionally important regions of individual repeats supports a hypothesis of concerted evolution and functional constraints acting together to homogenize *Aar\_AcSp1* repeats. In addition, several terminal region variants per individual *Argiope* genome indicate multiple *Argiope* *AcSp1* loci. Across *AcSp1* loci within an individual, we found homogenization of the repetitive region, but variation at the terminal coding regions. We also found evidence for stronger purifying selection in the N-terminal region versus the repetitive or C-terminal region, suggesting that the N-terminal region is the most constrained portion of the aciniform spidroin. The maintenance of multiple copies of *AcSp1* in *Argiope* genomes underscores the importance of aciniform silk in *Argiope* ecology and evolution. Indeed, variation between individuals and multiple gene copies within individuals could provide a method for the rapid synthesis of aciniform silk in this genus, and may represent the early stages of the differentiation that led to the extraordinary sequence and functional diversity of spider silks.

## Methods

### Isolation and sequencing of *AcSp1* containing BAC clone

A bacterial artificial chromosome (BAC) library was constructed by Rx BioSciences (Rockville, MD) with *Argiope argentata* genomic DNA inserted into pCC1BAC vector (Epicentre, Madison, WI). Colony pools were PCR screened for *AcSp1* with primers designed from the repetitive region of *Argiope trifasciata* *AcSp1* (Additional file 2: Table S1), resulting in one positive clone. The positive clone was restriction enzyme digested and a ~17 kb Hind III fragment of the full insert was found to contain the complete *AcSp1* gene.

The 17 kb fragment was gel purified with the S.N.A.P. UV-Free Gel Purification Kit (Invitrogen, Carlsbad, CA), ligated into HindIII digested pZerO™-2 plasmid (Invitrogen), and transformed into TOP10 cells (Invitrogen). Seven plasmid clones with the expected insert size and restriction enzyme digest patterns were end-sequenced with M13 and Sp6 primers to identify orientation of the inserts. Two clones (one of each insert orientation) were triple-digested with SpeI, XbaI, and XhoI and the fragments were gel-purified. The two clones were also single-digested with BamHI and the largest fragment from

each digest (5.5 kb or 5.9 kb, composed of the vector and either a 2.2 kb or 2.6 kb insert fragment) was gel-purified and re-circularized to produce subclones. The triple digest produced a 12.4 kb SpeI/XbaI fragment that was gel-purified and subcloned into SpeI digested pZerO-2 plasmid. End-sequencing the subclones revealed that the 2.6, 12.4, and 2.2 kb inserts corresponded to the *AcSp1* N-terminal, repetitive, and C-terminal encoding regions, respectively. The 2.6 and 2.2 kb fragments were sequenced in their entirety using primer walking (Additional file 2: Table S1). Because the 12.4 kb fragment contained repetitive nucleotide sequence, primer walking was not feasible. Instead, the 12.4 kb fragment was bidirectionally sequenced in its entirety using the transposon-based EZ-Tn5 < TET-1 > Insertion Kit (Epicentre Biotechnologies). The complete contig of the 17 kb genomic fragment was manually assembled with Sequencher 4.5 (Gene Codes, Ann Arbor, MI). An additional 1 kb of genomic sequence immediately adjacent to the 3' end of the 17 kb fragment was determined by primer walking (Additional file 2: Table S1) using the original BAC clone as template DNA. The complete *Argiope argentata* *AcSp1* gene was uploaded to GenBank with the accession number KJ206620.

### Inter- and Intraspecific sampling of N-, repetitive, and C-terminal coding region fragments

Genomic DNA was extracted from single legs removed from four *A. argentata*, one *A. aurantia*, and two *A. trifasciata* individuals using the DNeasy Blood & Tissue Kit (Qiagen, Valencia, CA). N-terminal, repetitive, and C-terminal encoding fragments of *AcSp1* were PCR amplified using primers designed from the *A. argentata* *AcSp1* complete gene (Additional file 2: Table S1).

PCR products of the expected size were purified using the AccuPrep Gel Purification Kit (Bioneer Inc., Alameda, CA). Products were directly sequenced. If a chromatograph had overlapping peaks, indicative of heterogeneous amplification, then the product was ligated into pJET 1.2 plasmid (ThermoScientific) and transformed into TOP10 cells. Individual colonies were PCR amplified using pJET1.2 Forward and Reverse sequencing primers. Inserts of the expected size were gel purified and sequenced. If one variant was highly abundant, then additional colonies were PCR amplified and digested with restriction enzymes to identify the abundant variants. The remaining undigested PCR products containing the rare variant were purified and sequenced.

### Diagnosing variants

Nucleotide sequences from the PCR fragments from each species were aligned as described below. For variant diagnosis, single nucleotide polymorphisms (SNPs) that were present in only one individual clone were

attributed to *Taq* polymerase error and that SNP was ignored. If a sequence had a pattern of polymorphism that was not present in at least one other clone, the sequence was discarded. Neighbor joining trees were then used to visualize highly similar sequences. Clusters that had greater than 95% identical sites were considered a variant group. With the exception of the cluster for *A. trifasciata* C-terminal coding variant 14 (95.2% identical sites), all clusters had greater than 98% identical bases. Clustered sequences were extracted and aligned to derive the majority rule consensus for that variant. Each variant is therefore supported by at least two sequences. Variants were uploaded to GenBank with accession numbers KJ206570–KJ206619.

### Phylogenetic analyses

The conserved spidroin N- and C-terminal regions from the complete *A. argentata* AcSp1 were aligned to 29 published spidroins that also have both N- and C-terminal regions (accession numbers in Additional file 2: Table S2) using ClustalW [49] implemented in Geneious v6.1.6 (Biomatters Ltd., Auckland, NZ). The N- and C-terminal regions were separately aligned with default settings and the alignments were adjusted by eye. The aa alignments dictated nucleotide alignments. N- and C-terminal encoding region alignments were concatenated for phylogenetic analyses of spidroin paralogs (Additional file 1: Figure S2). Despite potential recombination and convergence in the N- and C-terminal encoding regions, previous research found no conflict between the strongly supported nodes between separate N- and C-terminal trees and that concatenation of the terminal regions provides greater evolutionary resolution [12].

The 20 repeat units from the complete *A. argentata* AcSp1 repetitive region were divided into individual files and aligned as above with individual repeat units from published araneid AcSp1 sequences (Additional file 1: Figure S3, accession numbers in Additional file 2: Table S3). Alignments for the N- and C-terminal encoding sequences obtained from the PCR survey of individual genomes were created as above using diagnosed variants. Repetitive region alignments from the PCR survey also used the above method (alignments in Additional file 1: Figures S4–S6).

For each nucleotide alignment, bootstrap and maximum likelihood (ML) searches for optimal trees were simultaneously conducted over 5,000 replicates using RAXML 7.2.8 with the GTRGAMMA model [50,51] through the CIPRES webserver [52]. As implemented through CIPRES, RAXML has two substitution models: GTRGAMMA and GTRCAT. GTRGAMMA is considered more thorough [50–52]. Accession numbers given in Additional file 2: Tables S2–S4.

### Selection analyses

Estimates of the number of nonsynonymous substitutions per nonsynonymous sites (dN) and synonymous substitutions per synonymous sites (dS) were produced for three AcSp1 nucleotide alignments using MEGA5 [53]: N-terminal encoding variants (Figure 4B; Additional file 1: Figure S4), iterated repeats (Figure 3; Additional file 1: Figure S3), and C-terminal encoding variants (Figure 4C; Additional file 1: Figure S5). CodonTest [54] implemented through the Datamonkey webserver [55,56] indicated the Felsenstein 1981 (F81) [57] model of codon substitution as the best-fit for all analyzed datasets. dN/dS ratios less than, equal to, or greater than 1 were interpreted as purifying selection, neutrality, or positive selection, respectively.

### Availability of supporting data

All sequences generated in this study are deposited in GenBank (KJ206570–KJ206620). Alignments used in ML analyses are available shown in the additional files. Alignments and the corresponding trees for this study are available at TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S15355>).

### Additional files

**Additional file 1: Figure S1.** Nucleotide alignment of the 20 repeat units (Aar\_R) from the complete *A. argentata* AcSp1 repetitive region. Alignment position numbers shown in increments of ten. Frame 1 translation is shown under nucleotide sequences. Alignment prepared with Geneious v6.1.6 (Biomatters Ltd., Auckland, NZ). **Figure S2.** Nucleotide alignment of 30 concatenated AcSp1 N- and C-terminal coding regions. Alignment position numbers shown in increments of ten. Frame 1 translation is shown under nucleotide sequences. Alignment positions 1–510 encompass the N-terminal coding region, 511–840 the C-terminal coding region. Alignment prepared with Geneious v6.1.6 (Biomatters Ltd., Auckland, NZ) and available on TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S15355>). Abbreviations and accession numbers in Additional file 2: Table S2. **Figure S3.** Nucleotide alignment of Araneidae AcSp1 iterated repeats. Alignment position numbers shown in increments of ten. Frame 1 translation is shown under nucleotide sequences. Alignment prepared with Geneious v6.1.6 (Biomatters Ltd., Auckland, NZ) and available on TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S15355>). Abbreviations and accession numbers in Additional file 2: Table S3. **Figure S4.** Nucleotide alignment of AcSp1 N-terminal coding variants from PCR survey of individual *Argiope* genomes. Nucleotide alignment in FASTA format. Alignment available on TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S15355>). Abbreviations: *A. argentata* (Aarg), *A. aurantia* (Aau), and *A. trifasciata* (At). **Figure S5.** Nucleotide alignment of AcSp1 C-terminal coding variants from PCR survey of individual *Argiope* genomes. Nucleotide alignment in FASTA format. Alignment available on TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S15355>). Abbreviations: *A. argentata* (Aarg), *A. aurantia* (Aau), *A. trifasciata* (At), and *Araneus ventricosus* (Av). **Figure S6.** Nucleotide alignment of AcSp1 repeat region from PCR survey of individual *Argiope* genomes. Nucleotide alignment in FASTA format. Alignment available on TreeBASE (<http://purl.org/phylo/treebase/phyloids/study/TB2:S15355>). Abbreviations: *A. argentata* (Aarg), *A. aurantia* (Aau), *A. trifasciata* (At), and *Araneus ventricosus* (Av).

**Additional file 2: Table S1.** Primers used for full-length Aar\_AcSp1 sequencing and targeted amplification of N-terminal, repetitive, and C-terminal coding regions. The name and sequence of primers designed for



primer walking during BAC clone sequencing and primers designed to amplify N-terminal, repetitive, and C-terminal regions of *Aar\_AcSp1* are given. **Table S2.** Accession numbers of spidroin sequences used for maximum likelihood analysis of terminal regions (Figure 2). Spidroin name is the name used in this manuscript. Species is the spider species that corresponds to the N- or C-terminal accession number. If a full length gene was used, only one accession number appears. **Table S3.** Accession numbers for sequences used in maximum likelihood analyses of iterated repeats (Figure 3). Spidroin name is the name used in this manuscript. Species is the spider species that corresponds to the N- or C-terminal accession number. **Table S4.** Accession numbers for *AcSp1* sequences generated in this study and used in maximum likelihood analyses of repeat region and N- and C-terminal encoding variants (Figure 4). GenBank abbreviations, species, and accession number are given. **Table S5.** Predicted amino acid composition and codon usage of the coding region of *Aar\_AcSp1*. The percentage *Aar\_AcSp1* composed of each amino acid and percentage of each codon used for each amino acid. **Table S6.** Overall and putative helical region pairwise identities of *A. trifasciata* consensus *AcSp1* repeat aligned to consensus *AcSp1* repeat subparts of *L. hesperus* and *U. diversus*. Consensus repeat sequences from each subpart of *AcSp1* repeats from *L. hesperus* and *U. diversus* were aligned to a consensus repeat from *A. trifasciata*. Overall pairwise percent identity and percent identity shown for each of six helical regions as predicted by Xu et al [32].

#### Abbreviations

aa: amino acid; BAC: bacterial artificial chromosome; bp: base pairs; cDNA: complementary deoxyribonucleic acid; C-terminal: carboxy terminal; dN: non-synonymous substitution rate; dS: synonymous substitution rate; kb: kilobases; kDa: kilodaltons; ML: maximum likelihood; MYA: million years ago; NMR: nuclear magnetic resonance; N-terminal: amino-terminal; PCR: polymerase chain reaction; rDNA: DNA sequence that codes for ribosomal RNA; rRNA: ribosomal ribonucleic acid; SNP: single nucleotide polymorphism.

#### Competing interests

The authors declare that they have no competing interests.

#### Authors' contributions

RCC collected and analyzed data and drafted the manuscript, YZ, JW, RA, and KA collected data. NA contributed to conception and design of the study. CYH conceived the study, collected data, and drafted the manuscript. All authors reviewed and approved the final manuscript.

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